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(1) Ajrikant: BIQCHEM PHARMA INC. 2550 Pantel Johnson Beslevard, Suita 600 Level, Quebeo HTT SM (CA) (7) Inventor: Dionne, Gervale S31 Beulevard des Frairles, Building 19 Laval, Quebes ri7V 1B7 (CA)

(4) Representative: Ritter, Staphen David et al Mathys & Equire 10 Flact Street Leadon EC4Y 1AY (GB)

(5) 1.3-Oxethiolene nucleoside analogues.

The invention relates to 1,3-contriblers intelegate analogues and their use in the treatment of virsi invention. More specifically, this invention relates to (-)-4-amino-5-fluoro-1-(2-hydrusymethyl-1,3-contriblers-5-yi)-(1H)-pyrintide-2-one and pharmaceutical acceptable derivatives and pharmaceutical formulabors thereof.

EP 0 528 253 A1

The present invention relates to rusisaside analogues and their use in medicine. More specifically the imvention is concerned with 1,3-exemples nucleoside analogues, phermaceutical formulations thereof and the use thereof in the treatment of viral infections.

The drily compound ourrently approved for the treatment of conditions caused by MIV to 3'-ozido-3'deoxyrnymidine (AZT, zidovudine, BW 600U). However this compound has a eignificent elde-offcet liability and thus either cennot be carp byed or, carce employed, may have to be withdrawn in a significant number of patients. There is in consequence a continuing need to provide compounds which are effective against HIV but with a concommitant eignificantly better therepautic index.

The eampound of formula (1)

18

18

35

is a recernic marture of the two enantiomers of formulae (I-1) and (I-2):

We have now found that, surprisingly, the (-)-anantiomer of the compound of formula (I) is much more active than the (+)-are momer, although both enantiomers show unexpectedly low cylindustry. There is thus provided in a first aspect of the invention the (-) (or leavorobutory) ananthomer of the compound of formula (I) and pharmecoutically acceptable derivatives thereof.

The (-)-enentiomer has the chamical name (-)-4-amino-5-fluoro-1-(2-hydroxymethyl-1,3-exathlolan-5-yl)-(1H)-pyrimidin-8-one (hereins/far compound (A)). This exanticates "as the stretchists attraction to the stretchists are compound (A)).

Preferably compound (A) is provided substantially free of the curresponding (*)-anandomer, that is to say formula (i-1). no more than about 6% www of the (+)- energiomer, more preferably no more than about 2%, and most preferably

By "a pnarmaceutically acceptable dedivative" is meant any pharmaceutically acceptable soft, selec, or east less than about 1% w/w is present. of such eater, of compound (A) or any other compound which, upon administration to the recipient, is capable of providing (directly or indirectly) compound (A) or an antivirally active metabolite or residue thereof.

It will be appropriated by those sided in the art that compound (A) may be modified to provide pharmscouldeavy acceptants derivatives thereof, at functional groups in bett the bases mainty and at the hydroxymethyl group of the unathrolane ring. Modification at an auch functional groups are included within the scope of the Invention. However, of perticular interest are phermacautically acceptable derivatives obtained by modification of the 2-nydroxymethy) group of the excitioisne ring.

Preferred severs of compound (A) include the compounds in which the hydrogen of the 2-hydroxymethyl group is replaced by an any function

in which the non-namonyl moloty R of the ester is selected from hydrogen, straight or branched chain skyl (e.g., methoxymetryl), arrivyl (e.g., benzyl), arrivyl, n-putyl), alkoxysikyl (e.g., methoxymetryl), arrivyl (e.g., benzyl), arrivyllkyl (e.g., phenoxymethyl), aryl (e.g., phenoxymethyl), aryl (e.g., phenoxyl optionally substituted by halogen, C₁₋₁ elkyl or C₁₋₁ elkoxy); subhonate sewers such as sixyl- or aralkylsusphonyl (e.g., methanssulphonyl); arring acid esters (e.g., L-valyl or L-solaucyl) and mono-, 41- or at phosphate cetters.

With regard to the above described setters, unless otherwise appointed, any alkyl mainty present artwards grouply contains 1 to 18 carbon stamp, per faultrly 1 to 4 serbon stamp. Any any implety present in such assists advantageously comprises a phanyl group.

In particular the extens may be a O_{i+1} alkyl extension an unsubstituted benzyl extension on a unitaryl extensional distribution of logine), C_{1-4} alkyl, C_{3-4} alkowy, note or will unremetted orders.

Pharmaceutically acceptable salis of the compound (A) Include those derived from pharmaceutically acceptable inorganic and organic solds and beass. Examples of suitable edits include hydrochtoric, inviribromic, sulphuric hitris, perchloric, humaric, maleto, phosphoric, giveolic, isodic, editoria, succinic, toruser-o-sulphonic, tertario, edetic, cario, mathematulphonic, formic, benzoio, malonic, naphthalene-2-sulphonic and benzanasulphonic acids. Other acids such as oxalic, while not in thomselves pharmaceutically acceptable, may be useful as incorrectable in optaining the compounds of the invention and their pharmaceutically acceptable acid addition salits.

Saits derived from appropriate bases include alkali metal (e.g., application earth metal (e.g., magnesium), armenium and MR₂+ (where R is Q₁₋₄ alkyl) seits.

References terminafter to a compound according to the invention include both the compound (A) and the pharmaceutically acceptable districtives.

The community of the invention either transcrives possess antiviral activity and/or are metabolizable to such compounds, in particular these curricularies are effective in inhibiting the representation of retroviruses, including human retroviruses such as human immunodefficiency viruses (HIV's), the causetive agents of AIDS.

The compounds of the Invention are also useful in the treatment of enimals including man infected with the hepatitie B virus (HBV).

34

There is thus provided as a further aspect of the invention compound (A) or a pharmaceutically acceptable narrowive thereof for use as an active thereposite agent in particular as an activitie agent, for example in the year-next of resourcest infections or HBV infections.

In a further or alternative expect there is provided a method for the treatment of a viral infection, in particular with infection remained by HBV or a recrovirus suon as HIV, in a mammal including man comprising administration of an effective amount of compound (A) or a pharmacountricity acceptable derivative thereof.

There is also provided in a further or alternative expect use of compound (A) or a pharmaceutically appealed derivative thereof for the manufacture of a modicament for the treatment of a viral infestion.

The compounds of the invention are also useful in the treatment of AIDS related conditions such as AIDS-related complex (ARC), progressive generalized lymphedenopathy (PGL), AIDS-related neurological conditions (such a dementia or tropical paraparests), and HIV antibody positive and HIV-positive conditions. Kaposfa serooms, thrombodycoponia purpures and associated opportunistic infections for example preumocystic usurali.

The compounds of the invention are also useful in the prevention of progression to clinical linears of individuals who are enti-trity antibody or HIY-antigen positive and in prophylade following exposure to HIV.

The compound (A) or pharmaceutically occupable derivatives thereof they also be used for the prevention of viral contamination of physiological fluids such as blood or semen in vitro.

The compounds of the invention are also useful in the treatment of entirials limitaling man infected with the hepatitis 8 virus.

It will be appreciated by those skilled in the ert that reference heads to treatment extends to prophylade as well as the treatment of established infections or symptoms.

It will be further appreciated that the amount of a commount of the invention required for use in treatment will vary not only with the particular compound selected but also with the route of administration, the nature of the condition being wested and the age and condition of the patient and will be diffractely at the discretion of the attendant physician or vetorinarian, in general however a suitable dose will be in the range of from about 0.1 to about 750 mg/kg or bodywoight per day preferably in the range of 0.0 to do mg/kg/day, most preferably

EP 0 SES 288 A1

in the range of 1 to 20 mg/kg/4 to

15

The desired dose may conveniently be presented in a strictle dose or as divided dears administered at approprieto intervete, for example as evo, enrea tour or mere out doses per day.

The compound is conveniently administered in unit decage form; for example containing 10 to 1800 mg. conveniently 80 to 1000 mg, most conveniently 60 to 700 mg at active negrections per unit dossge form.

ideally the active ingredient should be seministered to achieve peak plasms concentrations of the active compound of frem about 1 to about 78 µM, preferably about 2 to 50 µM. most preferably about 3 to about 30 MM. This may be achieved, for example, by the intervenous injection of a 0.1 to 5% solution of the active ingradient, optionally in settine, or orally administered as a value containing about 1 to about 100 mg of the active ingradient. Desirable blood levels may be maintained by a continuous infusion to provide about 0.01 to shout 5.0 mg/kg/hour or by intermittent infusions containing about 0.4 to about 15 nig/kg of the active ingredient.

While it is possible that, for use in therapy, a compound of the invention may be administered as the rew chemical it is preferable to present the active ingredient as a pharmacoutical formulation.

The invention thus further provides a pharmaceutically formulation comprising compound (A) or a pharmaceutobily ecosphible derivative thereof together with one or more plantaneoutically acceptable cerriors memor and, optionally, other therapactic and/or crophylectic ingredients. The earner(s) must be 'sesspistie in the sense of being competitie will the uther ingredients of the formulation and not detected us to the

Phermacoutical formulations instude these suitable for trail rectal nase), (optical (including buccal and submedicated the remode. tingual), vaginal or parenteral (including intrarauscular, aub-cutaneous and intravenous) administration or in a form suitable for mindristration by infralation or insufficient. The formulations may, where sopropriate, be conveniently presented in discrete doesge units and may be prepared by any of the methods well known in the art of plasmacy. All methods hidded the step of bringing into association the active compound with liquid carriers or likely divided solid carriers or both and then, if necessary, shaping the product into the desired formulation.

Pharmaceutical formulations suitable for orei administration may conveniently be presented as discrete units such as capsules, cathets or tables each containing a predetermined amount of the active ingredient; as a powder or granutes; as a solution, a suspension of as an emulsion, it we solve ingredient may also be prosented as a bolue, electuary or pasts. Tablets and copoulos for ord administration may contain conventional exciplents such as pinding egents, fillers, lubriodists, disintegrants, of wetting agents. The tablets may be coated ecoording to methods well known in the art. One liquid proparations may be in the form of, for example, equadus or only suspensions, solutions, simulations, syrups or stixins, or may be presented as a dry product for constitution with water or other suitable vehicle before use. Buch liquid preparations may contain conventional additives such as suspending agents, emulativing agents, non-equeous vehicles (which may include edible of s), or pres-

Ine compounds eccording to the invention may also be formulated for parenteral administration (e.g., by grystives. injuction, for example below injection or continuous infusion) and may be presented in unit dose form in amposice, pre-filed syringes, small volume infusion or in multi-dose containers with an endan preservative. The compositions may take such forms as suspansions, solutions, or smulsions in only or equeous validates, and may contain formulatory agents such as suspending, stabilizing statics dispossing agents. Alternatively, the ective ingredient may be in powder form, obtained by accepte includen or storile exité or by lyuphilization from solution, for constitution with a suitable vehicle, e.g., sterile, pyrogen-free water, before use.

For topical administration to the epidernile the occupating according to the invention may be formulated se cintments, cesame or oxiona, or as a transdarmal natch. Cintments and creams may, for example, be formulated with an equecus or olly base with the addition of suitable thickening and/or goting agents, Lotions may be torsivisted with an equation or only base and will in general size contain one or more amulallying agents, subliking agents, dispersing agents, suspending agents, stockening agents, or exioting agents.

Formulations matable for topical administration in the mouth include lexenges comprising active ingredient in a flavored base, usually success and suscise or tragactanth; passiles comprising the active ingledient in an nert have such as galatin and glycerin or sucross and acacis; and mouthwasnes comprising the souve ingre

Phermaceutical formulations suitable for rectal administration wherein the carrier is a solid are most precioni in a sullative aquiti carrier. terably presented as unit dose suppositorise. Suitable carriers include cocca butter and other meterials commonty used in the art, and the suppositories may be conveniently formed by admissure of the active compound with the softened or melted carrier(s) tollowed by challing and shaping in moulds.

Formulations suitable for vaginal administration may be presented as pessaries, tempora, croams, gets, pusites, fusing or apraya containing in addition to the active ingredient such partiers as are known in the art to

Fur interment summitted due the compounds of the invention may be used as a liquid spray or dispessible

include rOR where R is an atiqui group, e.g., a C., atklyt group such se metry; or R is an acyl group, e.g., a C., atklyt group such as social or helogen, for example location, bromise or chiprine.

I he compound of formula (VIII) is conveniently reacted with 5-fillutro-cytosine or an appropriate cyrimiting boso produces the root (proviously saylated with a saylating agent such on hexamility distances) in a compatible privant such as methylene chloride using a Lewis sold such as thanking tetrachloride, trimethylally, triffeds, simethylally triffeds, atmostyrally locate (TMSI) or the (NY) compound such as SnCl.

The 1,3-executationes of formula (VIII) may be prepared for example by reaction of an eldehyde of formula (VII) with a merceptoscelal of formula (VI) in a compatible organic solvent, such as columns in the presence of an acid catagost for example a Lewis acid such as zinc chlorids.

The mercaptoscotals of formula (VI) may be prepared by auditicals known in the art. for exempte G. Hesse and L. Inster, Chem. Ser., 85, pp. 874-937 (1952).

The aldehydes of formule (VII) may be prepared by instructs brown in the set for example C.G. Halloquist and H. Hibbert, Can J. Resparch, 8, pp. 129-138 (1933). Conveniently the crude aldehyde (VII) may be putified by conveniently to the displacement of the displacement of the displacement of the displacement of the free aldehyde. In a second process the compound (A) is estained by base interconversion of a compound of formula (IX).

where \$ is a base convertible to \$-fluoro-cytosine. Such interconversion may be effected either by simple chemical transformation (e.g. the conversion of until) base to cytosine) or by an anzymatic conversion using a decryptionary transferace. Such methods and conditions for bear interconversion are well known in the art of mucleoside chemistry.

in a third precess a compound of fermula (XI)

94

may be converted to the compound (A) by conversion of the enonward NM_2 group to the S-flucibusine base by methods wall known in the nucleoside chemistry and

Many of the reactions described hereinabove have been extensively reported in the context of nucleoside systhesis, for example in <u>Nucleoside Analoss - Chemistry, biology and Madical Applications</u>, R.T. Walker real. Eds., Plentum Press, New York (1979) at pages 188-191 and T. Ueds. <u>Chymistry of Nucleosides and Nucleosides and Nucleosides.</u> Vol.1, L.B. Townsand Ed., Plentum Press, New York (1988) at pages 188-191, the discipance of which are incorporated by reference harebs.

It will be appreciated that the above reactions may require the line of, or conveniently may be applied to starting metadals having protected functional groups, and deprotection might thus be required as an intermediate or final aten to yield the cleared compared. Protection and deprotection of functional groups may be effected using currently as a protected from smally (e.g. benegit), soyl, anyl (e.g. 2,4-dintrophenyl) or skyl; subsequent removal of the protecting group being affected when dealted by hydrolysis or hydrogenolysis as appropriate using standard conditions. Hydroxyl chapter may be protected using any conventional mydroxyl protecting group, for example, as described in Property Groups in Organic Chemetry, J.P.W. McCente, Ed., Planum Press, New York (1973) or T.W. Greens, Protected Groups in Organic Chemetry, J.P.W. McCente, Ed., Planum Press, New York (1973) or T.W. Greens, Protected Groups in Organic Chemetry, John Wiley and Sons, New York (1984), Examples of subside hydroxyl protecting groups include groups selected from sityl (e.g., mothyl, t-buryl or methoxymetryl), erailyl (e.g., benzyl), and sityl groups such as trialkylskyl (e.g., t-butyldimetrylsdyl). The hydroxyl protecting groups may be removed by conveniental techniques. Thus, for example, sityl, skyl, scyl and heterocyclic groups may be removed by conveniental techniques. Thus, for example, sityl, skyl, and and heterocyclic groups may be removed by conveniental techniques. Thus, for example, sityl, skyl, and and heterocyclic groups may be removed by conveniental techniques.

may similarly be removed by solvolysis, e.g., by hydrolysis under scidic conditions. Arellylig mups reum as here Lyl cray to cleaved for example by treatment with DF yetherate and scotte unnyunde followed by removes of scalable groups so formed at an appropriate stage in the synthesis. Billyl groups may stan conveniently be removed using a source of fluoride ions even as betra-n-butylammonium fluoride.

In the above processes compound (A) is generally obtained as a mixture of the dis and trans isomers of

which the die learner to the compound of interest.

These services may be separated by physical means, e.g., chromatography on effice get or by fractional crystall Leadon, either checky or on a suitable derivative thereof, e.g., sosteres (propered for example with scado arrhydride) followed, after separate, by conversion back to the parent product (e.g., by descelylation with me-

Pharmaceutically ecceptable salts of the compounds of the Invention may be prepared as described in US menolic ammental). Petant No. 4,363,114, the disclosure of which is incorporated by reference herein. Thus, for example, when it is desired to prepare an acid addition selt of compound (A) the product of any of the above precedures may so converted into a sak by treatment of the resulting free base with a suitable solo using convention methods. Pharmaceutically acceptable acid addition salts may be prepared by reading the free base with an appropriate acid cournally in the presence of a sunspire solvent such as an ester (e.g., ethyl sectate) or an alcohol (e.g., metranol, ethanol or teopropenol), inergenic basic selts may be prepared by reasting the parent compound with a susable same such as an elookol (e.g., methenol). Shermecoullodly screprship salts may size he prepered from ether salts, including other phorma social ecceptable salts, of the compound (A) using conven-

Compound (A) may be sorror tod into a pharmacautically ecceptable phosphete or other seler by reaction with a prosphory/sting agent, such as POCI, or a suitable esterifying agent, such as an acid halide of anhydrido, as appropriate. Ar ester or sait of compound (A) may be converted to the parent compound for example

Resolution of the final product, or an intermediate or starting majorial therefor may be difected by any suitby hydrolysis. able method known in the art see for example E.L. Etel. Storeochemistry of Carbon Compounds, McGraw

Hall (1962) and S.H. Willen. Tables of Reserving Agents.

Thus for example the compound (A) may be obtained by ohirs! HPLC using a suitable estitionary phase for example scenyland proyeledentrin or cellulose triscetate and a suitable solvent for example an excelul such se otheror or an equeous solution of for example triathyl ammosium analais. Alternatively the corepreseds may be resolved by enzyme mediated enantional active catabolism with a suitable enzyme such as cylidina desmit ness or selective enzymetic degradation of a militable derivative a ff-ministrate. When selection is affected enzymetically the enzyme may be employed either in solution or, more conveniently, in immubilized form. Erzymas may be immobilized by any method known in the art, for example by adsorption onto a resin such as

The Invention will be further described by the following exemples which are not intended to limit the invention in any way. All temperatures are in degrees Colstus.

intermediate ?

25

(4)-0x-8-hydroxymothyi-8-(8'-fluorooytosin-1'-yi)-1,8-oxethiciane

() 2-Berzoyloxymethyl-5-acetony-1,3,nxetricishe

Benzpyloxysceusidehyde (216 33 g. 1.32 moi) was dissolved in pyridine (373 ml. 4.81 moi) and 1,4-dishlene -2,5-diol (100.31 g, 0.88 mel) was added to the solution. The hotorogenous mixture was stirred at 60-65°C under nüragen atmosphere for 1 hour. At the end of the reaction, a complete solution was obtained. Dichloromethane (860 ml) was added to the reaction mixture and it was ecoled to 0°0 with sait-les both. Acetyl chloride (251 ml, 3.95 mol) was added dropwise to the solution at 0-5°C over 1.5-2 hours. The reaction mixture was attreed at 0-5°C for 30 minutes, then it was poured corotally ento a cold (0°C) solution of saturated sodium be carbonate. The organic layer was separated. The water layer was extracted with dichloromathans (8 x 200 ml). The combined erganic layers were washed with esturated codium blearbonate sciution (5 x 200 ml) and brine (200 ml). The selution wee dried over sodium suitate and concentrated in vacua. The traces of pyridine were removed by executopic distillation with bonsono, \$20.79 g crude product was obtained which was purified by Kugatrohr distillation or filtration through a short allice sel ordumn. (Solvent system: hexane/ethyl sostate (3/1)).

(II) Cis-end trans-2-benzoylog methyl-5-(N, -acebd-5'-fluoro-ortosun-1'-vi)- 1.3-oxethiotene

5-Fluorocytosine (4.30 g. 33.5 mms), hexamethyldistazane (25 mi) and ammonium suthin (120 mg) were toiled under reflux unit the cytuative disserved (3 hours) and then further reflux district. The nexamethyldistazane was example to co-eveporate the activers. The resulting solution bis(tymethylatyl)-fluorocytosine in dichloromethane (40 ml) was acceed under argin to a solution of 2-bergoyloxymethyl -\$-acctany-1,3-oxatinotane (8.637 g. 30.3 mms) in dry dichloromethane (100 ml) and molecular sleves (44, 2 g) previously propared under argin and cooled at 0°C for 20 minutes. [(Trifluoromethane-suttony@coy) trimethyl allane (8 ml, 34 mms) was added to this mixture at 0°C and the resulting solution was strined at room temporature for 2 hours. The (fitties was chaken two times with 300 ml of brine and one time with distilled weter. The organic layer was dried over magnesium suffats, fittered and evaporated to dryriess. This afforded a brush 5-fluoro-cytosine derivative (10.1 g), Rr = 0.67 (EXDAC:MeOH 9:1).

This residue was acctyrated in the next step without further purification. The drude material W&& dissolved in dry dichieremethane (120 ml) in a 500 ml round bottom flask under argon. Triethylamine (12.7 ml, 91.1 mmol) and dimethyl aminopyridine (111 mg, 9.9 mmol) were added to the solution. The flask was their immersed in an ice bath for 1 hour under argon. Aceds anhydride (4.3 ml, 46 mmol), distilled over sodium scetate, was systemed attained social flask. The midure was stirred oversight and then carefully decented into an er-enmoyer flask containing seturated addium blosrbonate solution. The product was shon weathed with distilled water followed by brise softution. The methylene chicride portions were dried and evaporated under high vecoum to dryness, yielding an scetylated of picture as a color loss from weighing 9.8 g after drying. Flash ehromate graphy of the material using ethylecotate: mathenol (9.1) afforded 3.1 g. 7.8 mmol (46%) pure trans- and 3.5 g. 8.9 mmol (60%) pure efective compounds.

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trane-learner. R. = 0.65 in ethyl sostate/methanol 8:1
 U.V.: (MeOH) Lambda max: 909 nm
'H-NMR & (opm in COCL.)
 8.77 (b. 1H; Ca'-NH-Ao)
  8.06 (m. 2H: aromatic)
  7.70 (d. 1H; Cy'-15 Jos=8.3 Hz)
  7.82 (m. 1H; aromatio)
  7.49 (m, 21; aromatio)
  8.51 (dd, 1H; C-H)
  5.91 (dd. 1H; OyH)
  4.48 (dd, EH; OL-OH, OCOCHU)
  8.86 (dd, 1H; O4-H)
  7.24 (Ad. 1H. C.-H)
  2.86 (s, SH; NH-GOCHA)
  ris-incree: R. = 0.58 in ethyl acetate;methenoi 9:1
  U.V., (MarCH) Lambda max: 300 nm
1-NMR 8 (ppm in CDCL)
  8.72 (b, 1H; C.1-NH-AD)
  6.05 (m, 2H; aromatio)
  7 57 (d, 1H; Co'-H, Jos =6.2H2)
  7.80 (m. 1H; aromase)
  T.40 (m. 29t eromatic)
  6.32 (de, 1H) Ca+0
  6.47 (60, 1H; C+H)
  4 79 (44, 244 C<sub>1</sub>-CH_OCOC,H,)
  3.62 (60, 1Ht C.-H)
  3.19 (dd, 1Ht C. H)
  2.00 (8, 3H; NH-COCHL)
```

(III) (±)-U/s-hydroxymethyl-5-(5'-fluorocytosin-1'-VI) -1.3-oxethiolene

1.2 g (3.05 minor) of are 2-benzoyloxymetryl-d-(N₄-spetyl-d-fluorocytosts-1'-yl)-1,3-axishialane was attributed in 30 mi of metrenetic ammonia at 8°0 for 1 hour and their evernight at room temperature. The minutes was evaporated under reduced pressure. The residue was triturated twice (2 x 30 mil) with anhydrous either. The colid residue was recrystalitized in absolute otheriol to give 655 mg (2.84 mms), 87%) of pure c/s title prod-

MF & 536 366 A1

uct; m.D. 204-208°C; $R_{\rm f}$ = 0.21 in strykontain statement (6:1). The desired compound was identified by (H, (4C-NMR and U.Y. Lambda max (H₂O) 280.9 mm.

ale-learnes:

"H-NAR 8 (ppin in DM80-dg)
8.22 (d. 1H) Cg"-H, 1g; #7.26Hz)
7.84 (d. 2H; Cg"-NHg)
6.16 (l. 1H; Cg-H)
5.18 (l. 1H; Cg-CHg-OH)
5.18 (l. 1H; Cg-CHg-OH)
9.77 (m, 2H; Cg-CHg-OH)
9.36 (dd, 1H; Cg-H)
4-C-NMF (DM80-dg)

$$C_{6}'$$
 C_{2}' C_{4}' C_{5}'

153.46 158.14 134.63 126.32

 $(^{2}J_{CF} - 14.0H_{B})$ $(J_{CF} - 24.1H_{B})$ $(J_{CF} - 32.5H_{B})$

20 C_{4} C_{4} C_{2} C_{4} C_{5} C_{4} C_{5} C_{6} C_{6} 36.80 86.77 C_{6}

Example 1

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13

(+)-4-Amino-5-(luoro-1-(2-sydrony mothyl-1,5-oxethiolen-6-yl)-(1H)-pyrlinidis-2-une

() (-) Cla-3-hydraxymethyl-5-(5'-/fl. crossytuain-1'-yf)-1,3-uxatilislasia ironuphoaphata

To a stirred mixture of intermediate 1 (50-mg, 2,024 mixel) in dry timethyl phosphate (10 mi) cooled to O+C, was added drawins charpharus exychloride (1.22 mi, 13.1 mixel). The searcion mixture was attred at that immerature for 1 inc. 1 and their quenched in los water. The pH of the cold mixture was adjusted to 3 by the addition of arginalist 1N coldum hydroxide, then applied to a charcoal column (5 g, DARCO), which was clusted with water fulfored by otherior and equeous ammonts in a (10:10:1) ratio. Fractions containing crude of DEAE contains were combined and evaporated and subsequently was applied to a column containing 15 g of DEAE contained A29 (11CO₂-form). Elution was undertaken with a gradient or water (300 ml), 0.1M-NH₂HCO₃ (300 ml), and 0.2M NH₂HCO₃ (100 ml). Elution was undertaken with a gradient or water (300 ml), 0.1M-NH₂HCO₃ (300 ml), and 0.2M NH₂HCO₃ (100 ml). Evaporation of appropriate fractions effect dilution with water (90 ml) afforded (2) oth-2-hydroxymethyl-5-(6-fluoroxycosin-11-yl-1,8-catabiotens monophosphate as a wiste solid R₂ = 0.5 (0.0-CH:NH₂OH 644) yield = 812 mg, 1.77 mmol, 87.9%. 1H NMR 5 (ppm in D₂O), 8.27 (d, 1H, O₂H), J₃ = 6.47Hz), 6.33 (dd, 1H, O₂H), 5.47 (t, 1H, O₂H), 4.84 (m, 2H, O₂-CH₂OH), 3.63 (dd, 1H, C₂H3, 3.30 (dd, 1H, C₂H), HPLC>99%.

(ii) (+)-C/+2-hydroxymethyl-5-(6'-fluorocytodin-1' yl) ",8-exafhiciano

To a solution of (£) siz-2 hydroxymathyl-5-(6'-fluorocytosin-1'-yi)-1,3-exathiolane monophosphate (100 mg, 0.29 mmol) in 3 mi of glycine burier solution [glycine (52.6 mg) and magnesium chloride (19 mg) in water (10 mi)], was added in one portion 6'-nucleocidese [Sigme, 3.5 mg at 29 unit/mg). The resulting mixture was incubated st37°C with shaking. The reaction was monitored by HPLC (primal onlinen quantid glycoprotein (AGP) using 0.2M addism phosphate as elevant at pH ? with a flow rater 0.16 mixinting at different intervals. Only the (+)-onantiomer was observed after 2.5 hours. More enzyma (2 mg) was added, and insubstition was continued for a further 3 hours. HPLC analysis clearly showed selective and compliate hydrolysis of the (+)-arandomer. The resulting mixture was applied to a column of DEAE amphatiax 3-25 (HCO₀ form). Suiton was undertaken with violar (156 ml), followed by 0,1 and 0.2M Nit LiTCO₂ (100 ml each). Appropriate fresitive containing the first a user nucleocide were combined and concentrated. The remaining solid was purified on a short column allow using ethyl secrets, morehood (4.6:0.5) as aluent and than apparated by HPLC (employing the above mathronad conditions). This afforded ours (+)-ote-2-hydroxymathyl -5-(5'-fluorocytosis-1'-yl)-1,3-exathiolane (23 mg, 0.093 mmol, 32%) as a write solid (x)*131123°C (c, 1.00, MeOl (j m.p. 165°C NMR 6 (ppin in DMSC), 8.26

29 0 524 253 A1

(d, 1H, C_0 -H, $A_{t,t}$ =5.22 Hz), 7.87 (s, 1H, NH₂, D₂O exchangeable), 7.63 (s, 1H, NH₂, D₂O exchangeable), 6.20 (dd, 1H, C₀-H), 5.48 (t, 1H, C₂H), 6.24 (t, 1H, CH₂-QH, D₂O exchange), 5.84 (m, 2H, C₂-C_{H2}OH), 3.50 (dd, 1H, C₂H), 3.37 (dd, 1H, C₂H),

s (III) (-)- C/e-1-hydroxymethyl-5-(5'-fluorocytosin-1'-yl)-1,3-costhiolene

Appropriate fractions from the sephedax column containing the second sluted nucleoside described in step (ii) were combined and evaporated under reduced pressure. The residue was dissolved in 2 mi of water and rested with sikeline phosphetase (Sigma, 1 mi at 60 units/mi) followed by incubation at 37°C for 1.5 hours. Solvent was then evaporated and the residue was purified by column chromatography on stica get using EXACMEOH (4:1) as eluent followed by HPLC (separation using the same conditions mentioned above). This afforded pure (-)-cir-2-hydroxymethyi-5-(6'-fluoroxytosin-1'-yf)-1,3-coathotane (20 mg, 0.081 mmol, 28%) m.p., 180°C (d) rf=0.21. E:OAccMeOH (4:1), U.V.: (H₂O) max: 279.1nm. 'IH NMR 6 (ppm in DM8O-d₂), 8.16 (d. 1H, O'₂-H, J₁₀-7.26 Hz), 7.86 (b. 1H, O'₂-NH₂, D₂O exchangeable), 7.86 (b. 1H, C'₄-NH₂:D₂O exchangeable), 5.24 (t, 1H, C₂-H), 2.88 (m, 2H, C₂-C-12-O-H), 8.19 (dc, 1H, C₄-H), 3.16 (dd, 1H, C₄-H).

Intermediate 2 and Example 2 depict an alternate process for preparing the compound of formula (A).

Intermediate 2

(1'R, 2'8, 5'R)-MENTHYL-SR-(5'-FLUOROCYTISIN-1"-YL)-1,3-OXATHIOLANE-28-CARBOXYLATE

To a suspension of 5-filluprocytosine (165 mg. 1.2 mmol) in CPI-CI; (1 mL) at morn temperature under an argon atmosphere was added, successively, 2,4,8-collidine (0.317 mL, 2.4 mmol) and t-butyldimethylsilyl trifluoromethene-sulfonete (0.551 mL, 2.4 mmol). The resultant mixture was stirred for 16 minutes and a clear solution was obtained. A solution of (1'R,2'5,5'R)-menthyl-SR-acetoxy-1,3-oxathiclane-28-carboxylate (330 mg, 1 mmol) in CH₂Cl₂(0.5 mL) was introduced, followed by iodotrime tryistians (0.158 mL, 1.1 mmol). Stirring was continued for 3 hours. The mixture was diluted with CH₂Cl₂ (20 mL) and washed successively with seturated aqueous NaMSOs, water, brine and then was concentrated. The residue was taken up in either-haxanss (1:1, 10mL) and saturated equations NaHCO_c (2 mL) and stirred at room temperature for 15 minutes. The aquatum izyer was removed and the organic prisse was caratifuged to afford a while ledit within was washed with haxsince (3x5 mL) and then dried under vacuum. The product (1"R,2"5,5"R9-menthyt-5R-(5"-fluorobytosin-1"-yt)-1.3-examinisme-28-carboxylate (350 mg, 65%) thus obtained contained about 6% of (1/R.2/S.5/R)-membyl-68-(8"-fluorocytosin-1"-yr)-1.3-oxethiclene-25-carboxytate (NMR). This material was recrystalitized from MeCH/CH₂Ci/penzene to give a crystalline product: [c]₀²²+22* (a, 0.18, MeCH); m.g. 216-216*0. ¹H NMR (ODCh) 5 0.78 (d, SH. Je 7Hz), 0.91 (t, SH, Je7.8 Hz), 1.00 (m, 2H), 1.88-2.04 (m, 7H), 5.12 (dd, 1H, Je6.8 Hz, 8.1 Hz), 3.52 (dd, 1H, J=4.7 Hz, 8.1 Hz), 4.79 (dt, 1H, J=4.4 Hz, 4.8 Hz), 8.48 (R, 1 H), 8.76 (be, 1H, exchangeable), 6.42 (6t, 1H, J=5.0 Hz), 8.10 (bs. 1H, exchangentie), 8.48 (d. 1H, J=6.6 Hz); 190 NMR (CDCI_DMSO d.): 8 18.7, 21.2, 22.4, 23.7, 28.6, 31.8, 34.4, 38.6, 40.5, 47.2, 77.1, 79.1, 90.8, 128.3 (d. J=93 Hz), 137.1 (d. J=144 Hz), 184.2, 188.8 (d. J=18 Hz), 170.1.

Exemple 2

28-HYDROXYMFTHYL-SR-(5'-FLUOROCYTOBIN-1'-YL)-1,3-OXATHIOLANE

To a suspension of lithium aluminum hydride (10 mg, 0.54 mmol) in THF (1 mL) at ambient temperature under an argon atmosphere was alousy added a solution of (1/R.2/8,5/R)-menthyl-5R-(5"-fluorocytosin-1"-yl)-1.3-oxathiolane-28-carboxylate (54 mg, 0.135 mmol) in THF (2 mL). The seastion mixture was allowed to stir for 30 minutes, then quencined with excess methanol (2 mL), followed by the addition of allica gel (8 g). The resultant alumy was subjected to silica gel column chromatography (\$tOAo-Hexane-MeOH, 1:1:1) to provide a gummy solid which was dried exectropically with toluene to give 20.7 mg (#3%) of a white solid as the product: [a]p³⁶⁺114" (c, 0.12, MeOH); ¹H NMR (DMSO-d6) 8 3.14 (dd, 1H, J=6.8, 11.0 Hz), 3.42 (dd, 1H J=6.8, 11.9 Hz), 3.76 (m,2H), 5.16 (m, 1H), 5.42 (t, 1H, J=4.8 Hz), 6.14 (m, 1H), 7.50 (br m, 1H, exchangesbie), 7.83 (br m, 1H exchangesbie), 8.20 (d, 1H, J=7.66 Hz).

EP 0 000 255 A1

Exercis 1

Siciodical Astivity

(1) Antivirus Activity

Antiviral activity of the nompressed of Example 5 was determined against MEV-1 in the following call lines. C6160 cells, a human T-lymphoblastoid ost line, interest with rist-1 strain RF.

MT-4 cells, a human-T-cell leukaernia cell line, infected with HIV-1 strain RP.

Analytical activity in C6160 cells were determined by inhibition of synoytum formation (Tochikura et al Virology, 164, 542-648) and in MT-4 cells by inhibition of formazan conversion (Bobs et al. Biognam Biophys Res Commun., 142, pp. 128-134 (1987); Moseman, J.Immun. Meth., 59, pp. 65-67 (1983)]. Antiviral activities were also determined by analyzing the amount of HEV p24 antigen synthesized in the presence and absence of enantiomers.

The results are shown in Tables 1 and 2 below:

Table 1

20		50% Antiviral	Activity (µg/ml)
•	AZEAY	Pormanan	Inhibition of aynaytium formation
28	cells	HT-4	C8166
	Virus (HIV-1)	HIV-1 RF	MIV-1 RF
	(+)-enantiomer	> 1	0.04
30	(-)-enantiomer	0.14	0.0010
	Intermediate 1	0.065	0.013
**	AZT		0.0038

Table_2

sat.	Inhibition	HIV	p24	Synthosis	(µg/E1)
106	THRTALAM				

	Calls	, c	C\$166	
	Virus	3	JP	
13	(+) wanantiomer	d	.1	
	(-)-enantioner	(.0022	
w	Intermediate 1	(0.011	
₩	AZT	ŧ	0.017	

(II) Cytotoxicity

مد

The systems too of the compounds of Swample 1 and the recentle compound (Intermediate 1) were deserminas in two CD4 coti lines: #9 em CBM.

Compounds for test were serially diluted from 100 ug/mi to 0.3 µg/mi (final cancentrations) in 80 west mi-

EP 0 030 289 A1

crotitre plates. 3.8 x 10° calls were inoculated into each well of the plates including drug-free controls. After incubation at 37°C for 6 days, the viable cell count was determined by removing a sample of cell suspension and counting trypen blue excluding cells in a homocytometer.

The recuits are snown in Table 3.

Tabla 3

504	Cytotoxidity	140/=11

	Compound	CEH delle	H9 celle
14	(+) -enantioner	217	324
	(-)-enantioner	148	296
90	Intermediate 1	173	232

Claims

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- 1 (-)-d-emino-5-fluoro-1-(2-hydroxymethyl-1.3-oxathisten-5-y1)-(1H)-pydmidin-2-one or a pharmaceutholic celly acceptable derivative thereof.
 - 2. A compound according to claim 1 substantially free of the corresponding (+)-energlomer.
 - 3. A compound according to claim 1 wherein the (+)-enantiomer is present in an amount of no more than about 5% w/w.
 - A compound according to claim 1 wherein the (*)-anantiomer is present in an amount of no more than about 2% w/w.
 - A compound according to delm 1 wherein the (*)-ensulturier is present in an amount of less than about 1% w/w.
 - 6. A compound according to any proceeding claim in substantially pure form.
 - A charmaceutical composition comprising a compound according to any of claims 1 to 6 together with a pharmaceutically acceptable cerrier therefor.
 - 8. A compound according to any of claims 1 to 6 for use in therapy.
 - 9. Use of a compound according to any of claims 1 to 8 for the manufacture of a medicament for the treatment of a viral infection.
 - Use of 8 compound according to any of claims 1 to 6 for the manufacture of 8 medicament for the treatment of MIV intection.
 - Use of a compound according to any one of claims 1 to 8 for the manufacture of a medicament for the treatment of hepatitie 8 infection.
 - 12. A method for the preparation of a compound according to any of claims 1 to 8 which compilates asparations

HP 4 536 253 A1

of the (-)-ensatiomer from a mixture size containing the (+)-enantiomer.

- 13. A method aucording to plaim 12 wherein the mixture of compounds is a recessio mixture.
- 14. A method eccording to claim 12 or claim 15 wherein the separation is effected by chiral I:PLC.
 - 18. A method according to claim 14 wentein the MPLC employs as a stationary phrase acetylated \$- cyclo-dextrin or cellulose triacetate.
- 16. A method eccording to claim 12 or claim 13 wherein the separation is effected by enzyme-mediated enentioesective cetabolism.
 - 17. A method according to claim 16 wherein the enzyme is employed in immobilized form.
 - 18. A method according to claim 16 or claim 17 wherein the enzyme is cytidine desiminase.
 - 18. A method according to claim 18 or claim 17 wherein the enzyme is a 6'-nucleoligase.

Claims for the following Contracting Sistes : ES, GR

- 1. A method for the preparation of (-)-4-amino-5-fluoro-1-(2-hydroxymethyl-1,3-oxathiolan-5-y1)(1H)-pyrimidin-2-one or a pharmeceutically acceptable derivative thereof [compound (A)] which comprises the separation of the (-)-enantiomer from a mixture also containing the (+)-enantiomer.
 - A method eccording to claim 1 wherein compound (A) is obtained substantially free of the corresponding (+)-anantiomer.
 - A method according to claim 2 wherein the (+)-unentioned is present in an amount of sic more than about 0% w/w.
- 4. A mathed according to claim 2 wherein the (+)-enantiomer is present in an amount of no more than about 2% w/w.
 - 5. A method specialing to cisim 2 wherein the (+)-enentiaries is present in an amount of less than about 1% way.
 - 35 S. A method according to any precading claim wherein compound (A) to obtained in substantially pure form.
 - 7. A method according to any preceding claim wherein the muture of compounds is a recemb misture.
 - 5. A method according to any of claims t to 7 wherein the expersion is effected by chiral HPLC.
- A medical according to claim 6 wherein the MPLC employs as a stationary prime accordance proyacodecimal or cellulose tracetate.
 - 10. A method according to any one of delma 1 to 7 wherein the separation is effected by enzyme-mediated enterties elected by enzyme-mediated
 - 15. A method according to claim 10 wherein the enzyme is employed in immobilized form.
 - 12. A method according to claim 10 or claim 11 wherein the enzyme is cylidine desminase.
 - 13. A method according to claim 10 or claim 11 wherein the enzyme is a 5'-nucleotidate.
 - 14. A method for the preparation of a phermacautical formulation comprising as an active ingradient a compound produced according to any one of cisims 1 to 13 together with a phermacautically acceptable carrier therefor which method comprises admixture of the active ingradient and the carrier.

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